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# Minimizing the Impact of the Sample Matrix During Routine Pesticide Residue Analysis in Food

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#### **Abstract**

This application note describes the application of ultra sensitive detection to minimize the impact of matrix effects during analysis of 81 pesticide residues in a range of food products and also the use of a novel MRM acquisition mode that allows direct monitoring of the matrix background in each sample injected.

#### **Benefits**

- Detection of pesticides in complex food matrices using large multi-residue methods to below the required regulatory concentrations.
- · Ability to monitor changes in the sample matrix between samples and batches.
- · Reduction of matrix concentration to minimize matrix effects while maintaining detection.

#### Introduction

One of the biggest challenges in ensuring the safety of our food supplies is the measurement of hazardous ultra trace level components in the presence of a highly complex sample matrix. For the analysis of pesticides in food matrices, increased use of liquid chromatography systems, coupled with tandem quadrupole mass spectrometers has allowed progress in reducing the problems caused by the sample matrix. However, difficulties remain when trying to discriminate against matrix components that exhibit similar physiochemical properties. Unawareness of these difficulties in each unique sample can lead to poor quality results, and can impact a laboratory's performance and reputation.

Understanding the matrix challenge of each injected sample is clearly beneficial as is the ability to monitor changes in the sample matrix between samples and batches. This capability can lead to the continuous improvement of analytical quality in the laboratory. Conventional LC tandem quadrupole systems do not allow the direct monitoring of the sample matrix during high sensitivity MRM quantitation and it is only recently with the newest generation of instruments that this has become possible.

Problems caused by the sample matrix can include disruption to chromatography, increased chemical noise, and most notably, ionization suppression.<sup>1-4</sup> In highly complex matrices such as herbs and spices, these problems can be found in combination to make determination of pesticide residue concentration very difficult.

In addition to problems caused by the sample matrix, there are also pesticides that, by nature, are more difficult to analyze using LC-MS/MS due to a poor (relative) response factor. Successful analysis of these compounds to the regulatory concentration limits is difficult when considering the practicality of increasing sample amount and the balance of extracted matrix concentration. A much more practical solution is to use increased instrument sensitivity to maximize performance at these required concentrations. Also, if enough sensitivity is available, then the reduction of matrix concentration injected onto the system becomes possible.

Described here is the application of ultra sensitive detection to minimize the impact of matrix effects during analysis of 81 pesticide residues in a range of food products. Also described is the use of a novel MRM acquisition mode that allows direct monitoring of the matrix background in each sample injected.

#### Mass spectrometer acquisition

Quanpedia generated MRM parameters (a full MRM list can be found in Appendix 1) were used as the basis of RADAR-enabled mass spectrometer acquisition method. RADAR is an information-rich acquisition approach that allows measurement of target analytes with precision in MRM mode, while simultaneously scanning the background for all other components.

Figure 1 shows a RADAR-enabled mass spectrometer acquisition method with time scheduled MRMs for target pesticides and a simultaneous full scan (MS2) acquisition.

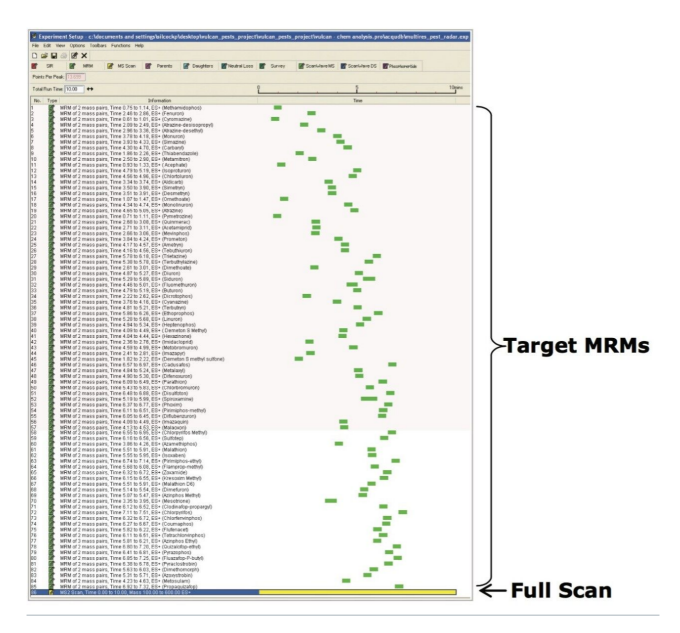


Figure 1. Mass spectrometer experiment showing RADAR acquisition mode.

# Experimental

Waters DisQuE (EN 15662:2008) Extraction Kit (QuEChERS) was used to prepare spiked extracts of grape, avocado, marjoram, and ginger. Sample matrix concentrations were 1g/mL for grape and avocado and 0.1 g/mL for marjoram and ginger. The final acetonitrile extracts from QuEChERS were diluted 10x into mobile phase and 10  $\mu$ L were injected onto the analytical system (referred to as original sample). Subsequent

dilutions of this were then made to reduce matrix effects.

#### LC conditions

LC system: ACQUITY UPLC

Column: ACQUITY BEH C<sub>18</sub> 100 mm x

2.1 mm, 1.7  $\mu m$ 

Mobile phase A: 0.1% HCOOH in H<sub>2</sub>O

Mobile phase B: 0.1% HCOOH in MeOH

Run time: 10.00 min

## UPLC gradient:

Time (min)	Flow (mL/Min)	%A	%B
-	0.5	90	10
0.25	0.5	90	10
7.75	0.5	2	98
8.5	0.5	2	98
8.51	0.5	90	10

#### MS conditions

MS system: Xevo TQ-S

Ionization mode: ES positive

Capillary voltage: 0.60 kV

Source temp: 130 °C

Desolvation temp: 650 °C

Cone gas flow: 150 L/hr

Desolvation gas flow: 1200 L/hr

### Results and Discussion

#### Detection to below regulatory limits

European Union (EU) regulations to control pesticide exposure from food consumption are among the toughest in the world. In order to import food and food commodities into Europe, the level of pesticide contamination must be below the stated maximum residue limits (MRLs) for that product.<sup>5</sup> Confirmation of positive results requires good quantitative performance well below these concentrations, which can be very challenging in more complex matrices.

Figure 2 shows a selection of extracted MRM chromatograms for pesticides spiked into avocado at 0.005 mg/kg. Quantitative and confirmatory transitions are both detected at this level, which is 10x below the European MRL (except zoxamide, which is 4x below). This includes parathion, which has a relatively poor response factor when analyzed using electrospray ionization. Comfortable quantitation of pesticides at these low concentrations allows high confidence when reporting results around maximum residue limits.

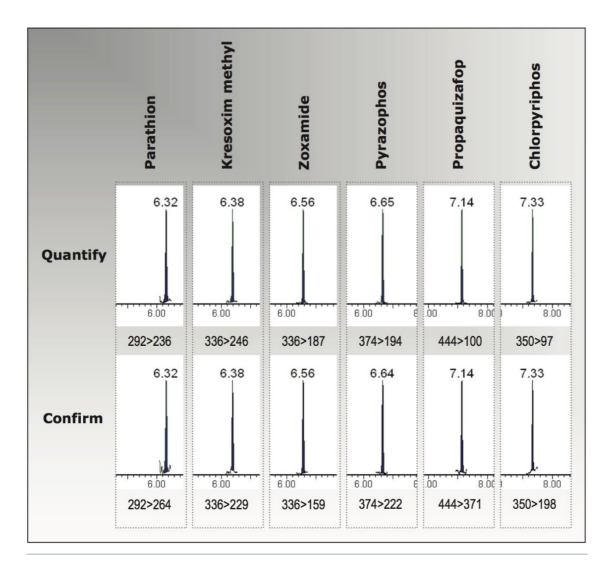


Figure 2. Quantitative and confirmatory MRM transitions for pesticides spiked into avocado at 0.005 mg/kg.

#### Monitoring matrix complexity

Each sample analyzed had full scan data available along with the MS/MS data. This was due to the RADAR functionality of the Xevo TQ-S being enabled. These data were used to monitor the complexity of the sample matrix background in each sample.

Differences in the co-extracted background for grape, avocado, marjoram, and ginger were observed by plotting the base peak intensity (BPI) chromatogram. For ginger and marjoram, 10x less sample was extracted using QuEChERS to give a 0.1 g/mL matrix, as opposed to the usual 1 g/mL matrix for grape and avocado. This is due to the extremely high complexity of the sample matrix, as well as to aid extraction of these drier samples. Figure 3 shows base peak intensity (BPI) chromatograms overlaid with MRM

chromatograms for pesticides spiked at  $1.0 \times 10^5$  g/kg for each matrix.

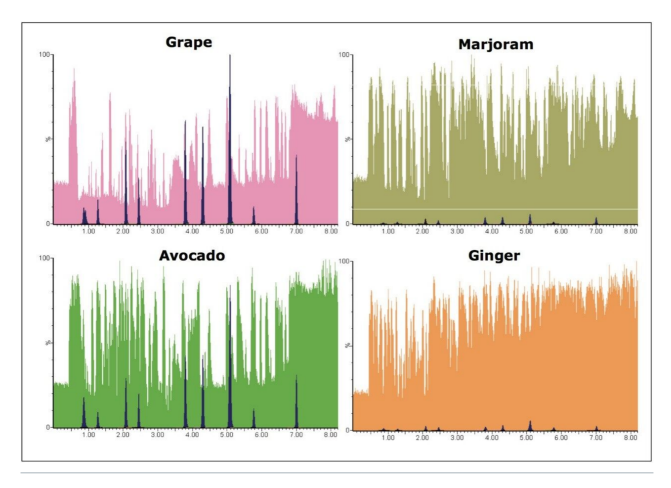


Figure 3. BPI chromatograms overlaid with MRM chromatograms for pesticides spiked at 0.01 mg/kg into grape (1.0 g/mL matrix), avocado (1.0 g/mL), marjoram (0.1 g/mL), and ginger (0.1 g/mL).

Despite the reduction in matrix concentration, the ionizable background is high in marjoram and ginger samples, compared with grape and avocado; as a consequence, the likelihood for analyte ion suppression (and enhancement) may be higher for these types of samples.

With simultaneous full scan it is also possible to observe specific components that co-elute with target analytes. Figure 4 shows BPI and MRM mass chromatograms for a grape sample spiked with dimethoate at 0.01 mg/kg. Full scan spectra from the elution region of dimethoate were combined and the most intense ion from the mass spectrum extracted into another chromatogram (XIC), revealing a discrete peak that co-elutes with dimethoate, as shown in Figure 4.

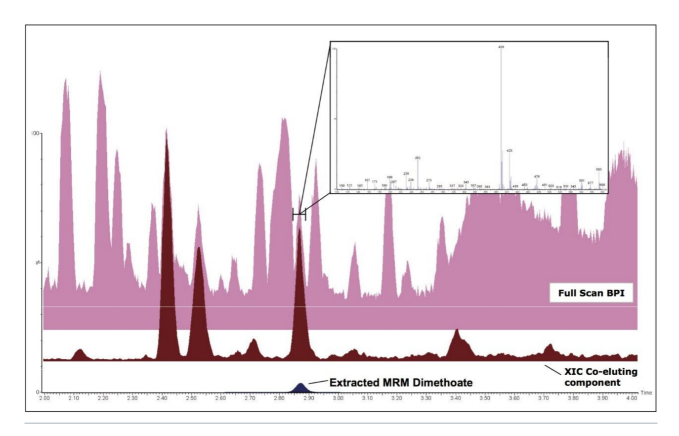


Figure 4. RADAR full scan BPI and MRM mass chromatograms for a grape sample spiked with dimethoate at 0.01 mg/kg. Also shown is the extracted ion chromatogram (XIC) of the co-eluting component with the subtracted mass spectrum inset.

If significant problems are observed with this or any other components in the matrix, the ability to observe them allows for further investigation and necessary remedial action to be carried out. Also, this acquisition mode can help to track the clean-up efficiency of the methodology employed.

#### Reduction of matrix effects

Minimizing matrix effects allows higher confidence in the quality of analytical data obtained. Reducing matrix concentration injected onto the analytical system is a simple and effective means to do this. When using a standard flow ESI source this can be achieved by reducing the amount of sample to be extracted, reducing the number of sample enrichment steps, or diluting final extracts. In any case, this is only a possibility if enough sensitivity is available to maintain detection at the required concentrations.

Ginger samples showed the highest ionizable background when compared to all other samples, despite having a relatively low matrix concentration (0.1 g/mL), as shown in Figure 3. Matrix effects were observed in the ginger samples with ion suppression and chromatography problems most apparent.

Diluting the ginger extracts 10x allowed recovery of distorted peak shape for cyromazine and reduction in matrix suppression for a number of pesticides, as shown in Figure 5. Table 1 shows reduction of ion suppression with a 10x dilution of sample. This reduction in suppression is clear when comparing peak area of pesticides in ginger to standards with no matrix present. As the matrix concentration is reduced the peak area response begins to correlate closely with standard peak areas.

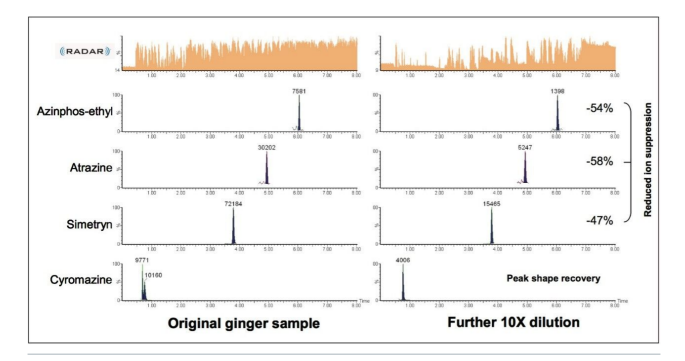


Figure 5. Effects of reducing sample matrix concentration by dilution for ginger. The full scan RADAR background is shown in the top chromatogram with MRM chromatograms for a selection of pesticides below.

	% Peak area recovery to standard			
	Original Extract	Diluted Extract		
Thiabendazole	89.2	105.2		
Atrazine-desisopropyl	71.6	100.8		
Aldicarb	36.4	91.2		
Desmetryn	49.2	97.0		
Prometon	85.8	109.2		
Simazine	63.1	103.4		
Hexazinone	80.0	98.7		
Demeton S Methyl	69.7	117.0		
Tebuthiuron	79.1	96.3		
Ametryn	66.7	103.4		
Terbutryn	81.7	102.8		
Azinphos Methyl	58.1	91.8		
Trietazine	46.8	91.6		
Azinphos Ethyl	60.5	86.1		

Table 1. Reduction of ion suppression for a ginger extract upon 10x dilution of original samples. Calculated as percent peak area recovery to a standard injection with no matrix present.

### Conclusion

- Xevo TQ-S allows detection of pesticides in complex food matrices using large multi-residue methods to below the required regulatory concentrations. This includes compounds with poor relative response factors.
- The RADAR mode of acquisition enables the collection of spectral information on background components in the sample matrix while simultaneously collecting MRM data. This can help identify areas of potential ion suppression, observe untargeted contaminants, and aid in the development of matrix reduction strategies.
- · Where matrix effects are observed, the high sensitivity offered by Xevo TQ-S allows matrix concentration in samples to be reduced to counteract these effects. This is possible while maintaining detection at regulatory concentrations and allows higher confidence in reported data.

# References

- 1. J M Marín *et al. Journal of Chromatography A*. 1216: 9, 1410-1420. 27 February 2009.
- 2. Hajšlová & Zrostlíková. *Journal of Chromatography A*. 1000:1-2, 181-197. 6 June 2003.
- 3. Gosetti *et al . Journal of Chromatography A*. 1217: 25, 3929-3937, 18 June 2010.
- 4. Kruve et al. Journal of Chromatography A. 1187: 1-2, 58-66, 11 April 2008.
- 5. website: http://ec.europa.eu/sanco\_pesticides/public/index.cfm

# Appendix 1 Pesticide MRM Parameters

	Precursor ion	Product ion	Collison (V)		Precursor ion	Product ion	Callison (V)
cephate	206 206	64 117	10 12	lmazapyr	262 262	69 86	24 24
cetamiprid	223	56	28	Imazaquin	312	86	26
ldicarb	223 213	126 89	12	Imidacloprid	312 256	267 175	18
CONCERTO	213 228	116 68	19 15	imiaaciopria	256 207	209 46	14
metryn	228	186	10	Isoproturon	207	72	20
trazine	216 216	96 174	34 16	Isoxaben	333 333	107 165	56 16
trazine-desethyl	188	79	21	Kresoxim Hethyl	336	229	15
SERVERY REST	188 174	146 79	17 25	200000000000000000000000000000000000000	336 249	246 160	15
trazine-desisopropyl	174	96	15	Linuron	249	182	15
zamethiphos	325 325	112 139	16 16	Malaoxon	315 315	99 127	22
zinphos Ethyl	368	132	22	Metalaxyl	280	192	16
-1.44.	368 340	160 132	35 15	W	280 203	220 104	12
zinphos Methyl	340	160	10	Metamitron	203	175	15
zaxystrobin	404 404	329 372	15 10	Methamidophos	142 142	94 125	12
uturon	237	84	28	Metabromuron	259	148	14
STORES CO.	237	126 131	14	Matandan	259 418	170	18 50
adusafos	271	159	28	Metosulam	418	175	26
arbaryl	202 202	117 145	20 15	Mevinphas	225 225	127 193	14
hlorbromuron	293	182	22	Monolinuran	215	99	32
y	293 350	204 97	12 15	***************************************	215 199	126 72	20 15
hlorpyrifos	350	198	20	Monuron	199	126	23
hlorpyrifas Methyl	322 322	125 290	25 15	Omethoate	214 214	125 183	20 10
hlortoluron	213 213	46	15 15	Parathion	292	236	12
lodina/op-propargyl	350	72 91	15	Phoxim	292 299	264 129	10
tournarop-proparysy.	350 363	266 289	16 30	Pissain	299 334	153 182	7 23
oumaphos	363	307	15	Pirimiphos-ethyl	334	198	21
yanazine	241 241	96 214	22	Pirimiphos-methyl	306 306	108	30 20
yromazine	167	60	23	Prometon	226	86	26
	167 253	108	15 17	20000000000000000000000000000000000000	226 444	184	16 15
emeton S Methyl	253	89	17	Propaguizafop	444	371	15
emeton S methyl sulfone	263 263	121 169	28	Pymetrozine	218 218	79 105	28 18
esmetryn	214	82	28	Pyraclostrobin	388	163	23
AND STATE OF THE S	214	172	19	TATA SERVICE STATE OF	388 374	194 194	30
icrotophos	238	193	10	Pyrazophos	374	222	20
ifenoxuron	287 287	72 123	18 18	Quinmerac	222 222	141 204	28 14
iflubenzuron	311	141	30	Quizalofop-ethyl	373	91	30
ACRES AND CONTRACT	311 339	158 72	15 24		373 233	299 94	16 23
imefuron	339	167	18	Siduron	233	137	15
imethoate	230 230	125 199	18	Simazine	202	96 124	22 16
imethomorph	388	165	28	Simetryn	214	96	23
ADD FOR A PORTUGUES	388 297	301 61	18 32	XXXXXII GARRET	214 298	124	18
isulfoton	297	89	12	Spiroxamine	298	144	19
luron	233 233	46 72	13 16	Sulfatep	323 323	97 171	30 14
thoprophos	243	97	29	Tebuthiuran	229	116	24
enuron	243 165	131 46	18	Terbuthylazine	229 230	172 96	16 26
	165 336	72	15 46	1000000 B000000	230 242	174 186	15 15
amprop-methyl	336	105	15	Terbutryn	242	200	15
uazafop-P-butyl	384 384	282 328	20 15	Tetrachlorvinghos	365 365	127 239	15 18
lufenacet	364	152	18	Thiabendazole	202	131	26
550000W00000	364 233	194 46	10 16	07-2507-10-10 07-2507-10-10	202 230	175 71	24 28
luomethuron	233	72	16	Trietazine	230	99	21
eptenophos	251 251	125 127	13 13	Zoxamide	336 336	159 187	36 23
lexazinone	253	71	28		] 330	101	

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